



INVESTOR IN PEOPLE

The Patent Office  
Concept House  
Cardiff Road  
Newport  
South Wales  
NP10 8QQ



I, the undersigned, being an officer duly authorised in accordance with Section 74(1) and (4) of the Deregulation & Contracting Out Act 1994, to sign and issue certificates on behalf of the Comptroller-General, hereby certify that annexed hereto is a true copy of the documents as originally filed in connection with the patent application identified therein.

In accordance with the Patents (Companies Re-registration) Rules 1982, if a company named in this certificate and any accompanying documents has re-registered under the Companies Act 1980 with the same name as that with which it was registered immediately before re-registration save for the substitution as, or inclusion as, the last part of the name of the words "public limited company" or their equivalents in Welsh, references to the name of the company in this certificate and any accompanying documents shall be treated as references to the name with which it is so re-registered.

In accordance with the rules, the words "public limited company" may be replaced by p.l.c., plc, P.L.C. or PLC.

Re-registration under the Companies Act does not constitute a new legal entity but merely subjects the company to certain additional company law rules.



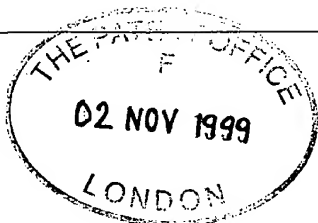
Signed *Andrews*

Dated 14 June 2000

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2 NOV 1999

Your reference  
PCS10370JWM-PROV

9925970.7

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#### Notes

Please type, or write in dark ink using CAPITAL letters. A prescribed fee is payable for a request for grant of a patent. For details, please contact the Patent Office (telephone 071-438 4700).

Rule 16 of the Patents Rules 1990 is the main rule governing the completion and filing of this form.

2 Do not give trading styles, for example, 'Trading as XYZ company', nationality or former names, for example, 'formerly (known as) ABC Ltd' as these are not required.

#### Warning

After an application for a Patent has been filed, the Comptroller of the Patent Office will consider whether publication or communication of the invention should be prohibited or restricted under Section 22 of the Patents Act 1977 and will inform the applicant if such prohibition or restriction is necessary. Applicants resident in the United Kingdom are also reminded that under Section 23, applications may not be filed abroad without written permission unless an application has been filed not less than 6 weeks previously in the United Kingdom for a patent for the same invention and either no direction prohibiting publication or communication has been given, or any such direction revoked.

The  
**Patent  
Office**

## Request for grant of a Patent Form 1/77

Patents Act 1977

### 1 Title of invention

1 Please give the title of the invention

TREATMENT OF PULMONARY  
HYPERTENSION

### 2 Applicant's details

☒ First or only applicant

2a If you are applying as a corporate body please give:

Corporate name  
PFIZER LIMITED

Country (and State of incorporation, if appropriate)  
UNITED KINGDOM

2b If you are applying as an individual or one of a partnership please give in full:

Surname

Forenames

2c In all cases, please give the following details:

Address  
RAMSGATE ROAD  
SANDWICH  
KENT

UK postcode CT13 9NJ  
(if applicable)

Country UNITED KINGDOM  
ADP number  
(if known)

689 2673 001

2d, 2e and 2f:

If there are further applicants  
please provide details on a separate  
sheet of paper.

☐ **Second applicant (if any)**

2d If you are applying as a corporate body please give:

Corporate name

Country (and State of incorporation, if appropriate)

2e If you are applying as an individual or one of a partnership please give in full:

Surname

Forenames

2f In all cases, please give the following details:

Address

UK postcode  
(if applicable)

Country

ADP number  
(if known)

3

An address for service in the United  
Kingdom must be supplied.

Please mark correct box

**3 Address for service details**

3a Have you appointed an agent to deal with your application?

Yes ☒ No ☐ ➡ go to 3b



*Please give details below*

Agent's name

J. W. MOORE

Agent's address

PFIZER LIMITED

RAMSGATE ROAD

SANDWICH

KENT

Postcode CT13 9NJ

Agent's ADP  
number

5629324001

*[Handwritten signature]*

3b:

If you have appointed an agent,  
all correspondence concerning  
your application will be sent to  
the agent's United Kingdom  
address.

3b If you have not appointed an agent please give a name and address in the United Kingdom to which all correspondence will be sent:

Name

Address

Postcode  
ADP number  
(if known)

Daytime telephone  
number (if available)

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## 5 Claiming an earlier application date

5 Are you claiming that this application be treated as having been filed on the date of filing of an earlier application?


Please mark correct box

Yes ☐ No ☒ **➡ go to 6**



***please give details below***

□ number of earlier application or patent number

 filing date

(day month year)

 and the Section of the Patents Act 1977 under which you are claiming:

Please mark correct box

15(4) (Divisional) ☐ 8(3) ☐ 12(6) ☐ 37(4) ☐

## 6

If you are declaring priority from a PCT Application please enter 'PCT' as the country and enter the country code (for example, GB) as part of the application number.

Please give the date in all number format, for example, 31/05/90 for 31 May 1990.

## 6 Declaration of priority

6 If you are declaring priority from previous application(s), please give:

Country of filing	Priority application number (if known)	Filing date (day, month, year)
1		

7

The answer must be 'No' if:  
 - any applicant is not an inventor  
 - there is an inventor who is not an applicant, or  
 - any applicant is a corporate body.

8

Please supply duplicates of claim(s), abstract, description and drawing(s).

Please mark correct box(es)

9

You or your appointed agent (see Rule 90 of the Patents Rules 1990) must sign this request.

Please sign here →

A completed fee sheet should preferably accompany the fee.

## 7 Inventorship

7 Are you (the applicant or applicants) the sole inventor or the joint inventors?

Please mark the correct box

Yes ☐ No ☒ →

A statement of Inventorship on Patents Form 7/77 will need to be filed (see Rule 15).

## 8 Checklist

8a Please fill in the number of sheets for each of the following types of document contained in this application.

Continuation sheets for this Patents Form 1/77

Claim(s)

Description

Abstract

Drawing(s)

8b Which of the following documents also accompanies the application?

Priority documents (please state how many)

Translation(s) of Priority documents (please state how many)

Patents Form 7/77 - Statement of Inventorship and Right to Grant (please state how many)

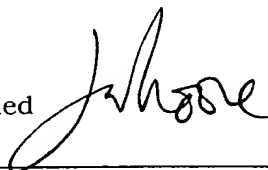
Patents Form 9/77 - Preliminary Examination/Search

Patents Form 10/77 - Request for Substantive Examination

## 9 Request

I/We request the grant of a patent on the basis of this application.

Signed



Date 01/11/1999

(day month year)

Please return the completed form, attachments and duplicates where requested, together with the prescribed fee to:

 The Comptroller  
 The Patent Office  
 Cardiff Road  
 NEWPORT  
 Gwent  
 NP9 1RH

The Comptroller  
 The Patent Office  
 25 Southampton Buildings  
 London  
 WC2A 1AY

TREATMENT OF PULMONARY HYPERTENSION

5        This invention relates to the use of the compound sildenafil, for the treatment of pulmonary hypertension.

      Pulmonary hypertension is a pathological condition in which the pulmonary artery pressure rises above normal levels and may cause sequelae of  
10    haemodynamic changes that can become life threatening. Symptoms of pulmonary hypertension include shortness of breath with minimal exertion, fatigue, dizzy spells and fainting. When pulmonary hypertension occurs in the absence of a known cause, it is referred to as primary pulmonary hypertension. Primary pulmonary hypertension is rare occurring in about 2 per million people worldwide.

15       Secondary pulmonary hypertension is much more common occurring as a result of other medical conditions, including congestive heart failure, chronic hypoxic lung disorder, including chronic obstructive pulmonary disease, inflammatory or collagen vascular diseases such as scleroderma and systemic lupus erythematosus,  
20    congenital heart diseases associated with left to right shunting and pulmonary thromboembolism.

      Sildenafil (Viagra<sup>®</sup>) is an orally-active, potent and selective inhibitor of cyclic guanosine monophosphate (cGMP) specific phosphodiesterase type 5 (PDE5) which  
25    is the predominant PDE isoenzyme in human corpora cavernosa. Consequently, it has been shown to be effective in the treatment of male erectile dysfunction. PDE5 is selectively abundant in the pulmonary vasculature compared to systemic vessels. Sildenafil increases intracellular concentrations of nitric oxide (NO) derived cGMP, thereby enhancing the effect of NO, and thus has potential to reverse metabolic and  
30    vascular defects in subjects with pulmonary hypertension.

It is known that inhaled NO stimulates the production of cGMP in pulmonary vascular smooth muscle cells resulting in selective pulmonary vasodilation.

Recently, it has been observed that administration of inhaled NO to subjects with

5 severe pulmonary hypertension resulted in significant decreases in pulmonary artery pressure and pulmonary vascular resistance without concomitant systemic hypotension. However the dose of inhaled NO is potentially limited by the formation of nitrogen dioxide, peroxynitrite or other toxic by-products. PDE5 is selectively abundant in the pulmonary vasculature in comparison to the systemic vessels and it  
10 has been observed that PDE5 is upregulated in pathological conditions leading to increase in pulmonary pressure. Sildenafil as a PDE5 inhibitor is expected to increase the level of cGMP and thus prolong the beneficial effect of the reduction of pulmonary blood pressure caused by NO with little effect on systemic blood pressure.

15

The use of phosphodiesterase inhibitors administered endotracheally or endobronchially (i.e. by inhalation) to treat pulmonary hypertension has been described in WO95/09636 but the compounds employed were neither particularly potent nor selective cGMP PDE inhibitors.

20

Sildenafil (5-[2-ethoxy-5-(4-methylpiperazin-1-ylsulfonyl)phenyl]-1,6-dihydro-1-methyl-3-propylpyrazolo[4,3-*d*]pyrimidin-7-one) and its preparation are described in European patent 0463756.

25

Thus according to the present invention we provide a method of treating a patient with pulmonary hypertension which comprises treating the patient with an effective amount of sildenafil or a pharmaceutical composition thereof.

30

The invention also provides for the use of sildenafil for the manufacture of a composition for treating pulmonary hypertension.



For use in the present invention sildenafil is preferably administered as a pharmaceutical composition. Thus, the compound can be administered in any conventional oral, parenteral, or transdermal dosage form, usually with a  
5 pharmaceutically acceptable carrier or diluent. Sildenafil is preferably employed in the form of its citrate salt but other pharmaceutically acceptable salts may also be used.

For oral administration a pharmaceutical composition can take the form of a  
10 solution, suspension, tablet, pill, capsule, powder or the like. Tablets containing various excipients such as sodium citrate, calcium carbonate and calcium phosphate are employed along with various disintegrants such as potato or tapioca starch and certain complex silicates, together with binding agents such as polyvinylpyrrolidone, sucrose, gelatin and acacia. Additionally, lubricating agents such as magnesium  
15 stearate, sodium lauryl sulfate and talc are often used for tableting purposes. Solid compositions of a similar type are also employed as fillers in soft and hard-filled gelatin capsules; preferred materials in this connection also include lactose or milk sugar as well as high molecular weight polyethylene glycols. When aqueous suspensions and/or elixirs are desired for oral administration, the compounds can be  
20 combined with various sweetening agents, flavoring agents, colouring agents, emulsifying agents and/or suspending agents, as well as such diluents as water, ethanol, propylene glycol, glycerin and various like combinations thereof.

For purposes of parenteral administration, solutions in sesame or peanut oil or  
25 in aqueous propylene glycol can be employed, as well as sterile aqueous solutions of the corresponding water-soluble salts. Such aqueous solutions may be suitably buffered, if necessary, and the liquid diluent first rendered isotonic with sufficient saline or glucose. These aqueous solutions are especially suitable for intravenous, intramuscular, subcutaneous and intraperitoneal injection purposes. In this  
30 connection, the sterile aqueous media employed are all readily obtainable by standard techniques well-known to those skilled in the art.

Sildenafil may also be administered as an inhaled formulation and this may have advantages in delivering the active compound directly to the lung area. Suitable formulations for inhaled administration are described in our co-pending British patent application number 911826.7.

Methods of preparing various pharmaceutical compositions with a certain amount of active ingredient are well known to those skilled in this art, or may be determined by reference to literature precedents.

10

The exact dose of sildenafil administered will, of course, differ depending on the subject being treated, on the severity of the condition, on the manner of administration and on the judgment of the prescribing physician. Thus, because of patient-to-patient variability, the dosages given below are a guideline only and the physician may adjust doses of the compounds to achieve the treatment that the physician considers appropriate for the patient. In considering the degree of treatment desired, the physician must balance a variety of factors such as the age of the patient and the presence of other diseases or conditions (e.g. cardiovascular disease). In general, the compound will be administered in a range of from 0.5 to 300 mg per day, preferably 5 to 125 mg per day, more preferably 25-100 mg per day.

20

Sildenafil may also be administered in conjunction with the administration of nitric oxide to treat pulmonary hypertension.

25

In addition to treatment of adult patients, a further application of the invention is in the treatment of very young children born with congenital heart disease. Sildenafil can be used to treat pulmonary hypertension in such subjects and can thus delay the immediate need for surgery until the patient is better able to withstand the trauma of surgery. Sildenafil can also be used to treat children who have pulmonary hypertension post operatively or due to respiratory distress syndrome or neonatal hypoxia.

30

CLAIMS

1. A method of treating a patient suffering from pulmonary hypertension which  
5 comprises treating said patient with an effective amount of sildenafil or a  
pharmaceutical composition thereof.
2. The use of sildenafil for the manufacture of a pharmaceutical composition for  
the treatment of pulmonary hypertension.

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